

=> s guanine nucleotide exchange factor
L1 4246 GUANINE NUCLEOTIDE EXCHANGE FACTOR

=> s rac
L2 17564 RAC

=> s gef

L3 2034 GEF

=> s exchange factor
L4 6154 EXCHANGE FACTOR

=> s l1 or l3 or l4
L5 6942 L1 OR L3 OR L4

=> s l2 and l5
L6 754 L2 AND L5

=> s polypeptide? or peptide? or protein?
2 FILES SEARCHED...
L7 6849438 POLYPEPTIDE? OR PEPTIDE? OR PROTEIN?

=> s l2 and l5 and l7

L8 704 L2 AND L5 AND L7

=> dup rem l8

PROCESSING COMPLETED FOR L8
L9 290 DUP REM L8 (414 DUPLICATES REMOVED)

=> s l9 and py<1998
1 FILES SEARCHED...
3 FILES SEARCHED...
4 FILES SEARCHED...
L10 46 L9 AND PY<1998

=> d l10 ibib abs 1-46

L10 ANSWER 1 OF 46 BIOSIS COPYRIGHT 2003 BIOLOGICAL
ABSTRACTS INC.
ACCESSION NUMBER: 1998:32642 BIOSIS
DOCUMENT NUMBER: PREV199800032642
TITLE: Cbl-b, a member of the Sli-1/c-Cbl ***protein***
family, inhibits Vav-mediated c-Jun N-terminal kinase
activation.
AUTHOR(S): Bustelo, Xose R. (1); Crespo, Piero; Lopez-Barahona,
Monica; Gutkind, J. Silvio; Barbacid, Mariano
CORPORATE SOURCE: (1) Dep. Mol. Oncol., Bristol-Myers Squibb
Pharm. Res.
Inst., Princeton, NJ 08543 USA
SOURCE: Oncogene, (***Nov. 20, 1997***) Vol. 15, No. 21, pp.
2511-2520.
ISSN: 0950-9232.

DOCUMENT TYPE: Article
LANGUAGE: English

AB We have used the yeast two-hybrid system to identify ***proteins***
that interact with Vav, a GDP/GTP ***exchange*** ***factor*** for
the ***Rac*** -1 GTPase that plays an important role in cell signaling
and oncogenic transformation. This experimental approach resulted in the
isolation of Cbl-b, a signal transduction molecule highly related to the
mammalian c-cbl proto-oncogene product and to the C. elegans Sli-1
protein, a negative regulator of the EGF-receptor-like Let23
protein. The interaction between Vav and Cbl-b requires the
entire

SH3-SH2-SH3 carboxy-terminal domain of Vav and a long stretch of
proline-rich sequences present in the central region of Cbl-b. Stimulation
of quiescent rodent fibroblasts with either epidermal or platelet-derived
growth factors induces an increased affinity of Vav for Cbl-b and results
in the subsequent formation of a Vav-dependent trimeric complex with the
ligand-stimulated tyrosine kinase receptors. During this process, Vav, but
not Cbl-b, becomes highly phosphorylated on tyrosine residues.
Overexpression of Cbl-b inhibits the signal transduction pathway of Vav
that leads to the stimulation of c-Jun N-terminal kinase. By contrast,
expression of truncated Cbl-b ***proteins*** and of missense mutants
analogous to those found in inactive Sli-1 ***proteins*** have no

detectable effect on Vav activity. These results indicate that Vav and
Cbl-b act coordinately in the first steps of tyrosine ***protein***
kinase receptor-mediated signaling and suggest that members of the
Sli-1/Cbl family are also negative regulators of signal transduction in
mammalian cells.

L10 ANSWER 2 OF 46 BIOSIS COPYRIGHT 2003 BIOLOGICAL
ABSTRACTS INC.
ACCESSION NUMBER: 1998:8424 BIOSIS
DOCUMENT NUMBER: PREV199800008424
TITLE: The ***guanine*** ***nucleotide*** ***exchange***

factor Tiam1 affects neuronal morphology: Opposing
roles for the small GTPases ***Rac*** and Rho.

AUTHOR(S): van Leeuwen, Frank N.; Kain, Hendrie E. T.; Van Der
Kammen,

Rob A.; Michiels, Frits; Kranenburg, Onno W.; Collard, John
G. (1)

CORPORATE SOURCE: (1) Netherlands Cancer Inst., Div. Cell Biol.,
Plesmanlaan

121, 1066 XC Amsterdam Netherlands

SOURCE: Journal of Cell Biology, (***Nov. 3, 1997***) Vol.
139,

No. 3, pp. 797-807.

ISSN: 0021-9525.

DOCUMENT TYPE: Article
LANGUAGE: English

AB The invasion-inducing T-lymphoma invasion and metastasis 1 (Tiam1)
protein functions as a ***guanine*** ***nucleotide***
exchange ***factor*** (***GEF***) for the small GTPase
Rac1. Differentiation-dependent expression of Tiam1 in the developing
brain suggests a role for this ***GEF*** and its effector Rac1 in the
control of neuronal morphology. Here we show that overexpression of
Tiam1

induces cell spreading and affects neurite outgrowth in N1E-115
neuroblastoma cells. These effects are ***Rac*** -dependent and
strongly promoted by laminin. Overexpression of Tiam1 recruits the
alpha6beta1 integrin, a laminin receptor, to specific adhesive contacts at
the cell periphery, which are different from focal contacts. Cells
overexpressing Tiam1 no longer respond to lysophosphatidic acid-induced
neurite retraction and cell rounding, processes mediated by Rho,
suggesting that Tiam1-induced activation of ***Rac*** antagonizes
Rho

signaling. This inhibition can be overcome by coexpression of
constitutively active RhoA, which may indicate that regulation occurs at
the level of Rho or upstream. Conversely, neurite formation induced by
Tiam1 or Rac1 is further promoted by inactivating Rho. These results
demonstrate that ***Rac*** - and Rho-mediated pathways oppose each
other during neurite formation and that a balance between these pathways
determines neuronal morphology. Furthermore, our data underscore the
potential role of Tiam1 as a specific regulator of ***Rac*** during
neurite formation and illustrate the importance of reciprocal interactions
between the cytoskeleton and the extracellular matrix during this process.

L10 ANSWER 3 OF 46 BIOSIS COPYRIGHT 2003 BIOLOGICAL
ABSTRACTS INC.
ACCESSION NUMBER: 1997:355832 BIOSIS
DOCUMENT NUMBER: PREV199799662235

TITLE: Identification of a novel, putative Rho-specific GDP/GTP
exchange ***factor*** and a Rho-A-binding
protein : Control of neuronal morphology.

AUTHOR(S): Gebbink, Martijn F. B. G.; Kranenburg, Onno; Poland,
Mieke;

Van Horck, Francis P. G.; Houssa, Brahim; Moolenaar, Wouter
H. (1)

CORPORATE SOURCE: (1) Div. Cell. Biochem., Neth. Cancer Inst.,
Plesmanlaan

121, 1066 CX Amsterdam Netherlands

SOURCE: Journal of Cell Biology, (1997) Vol. 137, No. 7, pp.
1603-1613.

ISSN: 0021-9525.

DOCUMENT TYPE: Article
LANGUAGE: English

AB The small GTP-binding ***protein*** Rho has been implicated in the
control of neuronal morphology. In N1E-115 neuronal cells, the
Rho-inactivating C3 toxin stimulates neurite outgrowth and prevents
actomyosin-based neurite retraction and cell rounding induced by
lysophosphatidic acid (LPA), sphingosine-1-phosphate, or thrombin acting

WEST Search History

10/054 435
A#8

DATE: Thursday, April 03, 2003

| <u>Set Name</u> side by side | <u>Query</u> | <u>Hit Count</u> | <u>Set Name</u> result set |
|--|-----------------------------------|------------------|-------------------------------|
| <i>DB=USPT,PGPB,JPAB,EPAB,DWPI; PLUR=YES; OP=ADJ</i> | | | |
| L8 | l1 with L7 | 292 | L8 |
| L7 | protein or polypeptide or peptide | 359298 | L7 |
| L6 | l1 with L5 | 35 | L6 |
| L5 | l2 or L4 | 628 | L5 |
| L4 | guanine nucleotide exchange | 362 | L4 |
| L3 | l1 with L2 | 23 | L3 |
| L2 | gef! | 369 | L2 |
| L1 | rac! | 4167 | L1 |

END OF SEARCH HISTORY